

A step toward red-light regulated optogenetic tools

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A schematic representation of the illumination of the sensor domain of a photoreceptor and the molecular propagation of the light signal to the effector (in red on the right-hand edge of the image). Credit: TU Graz/IBC

The aim of optogenetics is to control genetically modified cells using light. A team of Graz scientists led by Andreas Winkler from the Institute of Biochemistry at TU Graz have set a milestone in the future development of novel red-light regulated optogenetic tools for targeted cell stimulation. For the first time ever, they were able to observe



molecular principles of sensor-effector coupling in the full-length structure of a red-light responsive protein and describe detailed mechanisms of signal transmission over long distances at a molecular level. The results of the research have been published in the open access journal *Science Advances*.

Helix as light switch

To survive, cells and organisms have to adapt to new environmental conditions. This is the job of "protein building blocks" that interact with each other in different ways, thus creating cellular networks allowing adaptations to be made to changed environments. The sensors or "receptors" of external stimulation, such as light, are at least in part coupled to specific effectors in order to specifically activate or inhibit cellular signal molecules depending on need.

The Graz research group, which comprises not only researchers from TU Graz but also scientists from the Medical University of Graz, presents molecular details of a red-light photoreceptor involved in the production of a central bacterial messenger molecule, and describes the structure of a full-length light receptor together with its enzymatic effector for the first time. The architecture and composition of the linker element connecting the sensor and effector is very important in light regulation.

TU Graz biochemist Andreas Winkler, head of the research group, says, "By using a combination of X-ray structural analysis and hydrogendeuterium exchange by which the structural dynamics and conformational changes can be analysed, we managed to better understand the functional characteristics of this helical coupling element. We were able to show that illuminating the sensor with red light resulted in a rotation-like change in the coiled coil linker region, which in turn effects the enzymatic activity of the neighbouring effector." The Graz researchers were thus able to determine structural details of a red-light



regulated full-length system and describe molecular mechanisms of signal transduction.

Towards a rational design of proteins

The research contributes to better understanding the modularity of naturally occurring protein domains and being able to develop new optogenetic tools. Diverse combinations of different sensor modules are found in nature, such as red-light sensors, blue-light sensors and pH sensors—sometimes with identical and sometimes different effectors. From this, the researchers conclude that there are molecular similarities in signal transduction and therefore that rational and completely arbitrary combinations of sensors and effectors which do not occur in nature are conceivable.

Andreas Winkler says, "We are currently limited to naturally occurring systems to a great extent in the use of directly regulated enzymatic functionalities. The long-term aim is to generate new <u>light</u>-regulated systems which can overcome the limitations of nature and which would be of great interest for different applications in optogenetics."

More information: "Long-range allosteric signaling in red light–regulated diguanylyl cyclases" *Science Advances*, advances.sciencemag.org/content/3/3/e1602498

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